

## EAST Search History

| Ref # | Hits   | Search Query   | DBs                                       | Default Operator | Plurals | Time Stamp       |
|-------|--------|--|---|------------------|---------|------------------|
| L1    | 379368 | tolerance or tolerize or toleragenic   | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L2    | 335    | mHag or (minor adj histocompatibility adj antigen)   | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L3    | 254065 | transplant or transplantation or transplanted or transplanting or graft or grafted or grafting | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L4    | 177929 | reject or rejection  | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L5    | 100    | L1 and L2 and L3 and L4  | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L6    | 2371   | HA-1 or ha1 or (ha adj "1")  | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L7    | 379368 | tolerance or tolerize or toleragenic   | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L8    | 36942  | ((514/2) or (514/8) or (514/12) or (530/350) or (530/328) or (530/327)).CCLS.                  | US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT | OR               | OFF     | 2006/08/29 14:20 |
| L9    | 247    | L8 and L6  | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:20 |
| L10   | 105    | L9 and L7  | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:21 |
| L11   | 2371   | HA-1 or ha1 or (ha adj "1")  | US-PGPUB; USPAT; EPO; JPO; DERWENT        | ADJ              | ON      | 2006/08/29 14:21 |

## EAST Search History

|     |         |  |   |     |    |                  |
|-----|---------|--|---|-----|----|------------------|
| L12 | 12816   | GvHD or GvH or (graft adj versus<br>adj (host or leukemia)) or GvL   | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | ADJ | ON | 2006/08/29 14:21 |
| L13 | 6757711 | treat or treatment or prevent or<br>prevention or eliminate or<br>elimination or therapy or<br>therapeutic | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | ADJ | ON | 2006/08/29 14:20 |
| L14 | 11      | L11 and L1 and L2 and L12 and L3<br>and L4 and L13   | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | ADJ | ON | 2006/08/29 14:20 |
| L15 | 8250    | goulimy.in. or hunt.in. or englhard.<br>in.  | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | ADJ | ON | 2006/08/29 14:22 |
| L16 | 8536    | goulimy.in. or hunt.in. or<br>engelhard.in.  | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | ADJ | ON | 2006/08/29 14:22 |
| L17 | 311     | I16 and (I7 or I6 or I2)   | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | ADJ | ON | 2006/08/29 14:23 |
| L18 | 14      | I16 and I7 and ( I6 or I2)   | US-PGPUB;<br>USPAT;<br>EPO; JPO;<br>DERWENT | ADJ | ON | 2006/08/29 14:23 |

# SCORE Search Results Details for Application 10623176 and Search Result us-10-623-176a- 2.rag.

|                                 |   |                                       |                           |  |
|---------------------------------|---|---------------------------------------|---------------------------|--|
| <a href="#">Score Home Page</a> | <a href="#">Retrieve Application List</a> | <a href="#">SCORE System Overview</a> | <a href="#">SCORE FAQ</a> | <a href="#">Comments / Suggestions</a> |
|---------------------------------|---|---------------------------------------|---------------------------|--|

This page gives you Search Results detail for the Application 10623176 and Search Result us-10-623-176a-2.rag.

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GenCore version 5.1.9  
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OM protein - protein search, using sw model

Run on: August 18, 2006, 00:39:56 ; Search time 191 Seconds  
(without alignments)  
21.544 Million cell updates/sec

Title: US-10-623-176A-2

Perfect score: 45

Sequence: 1 VLHDDLLEA 9

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 2589679 seqs, 457216429 residues

Total number of hits satisfying chosen parameters: 442222

Minimum DB seq length: 0

Maximum DB seq length: 9

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : A\_Geneseq\_8:\*  
1: geneseqp1980s:\*  
2: geneseqp1990s:\*  
3: geneseqp2000s:\*  
4: geneseqp2001s:\*  
5: geneseqp2002s:\*  
6: geneseqp2003as:\*  
7: geneseqp2003bs:\*  
8: geneseqp2004s:\*  
9: geneseqp2005s:\*  
10: geneseqp2006s:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

[SUMMARIES](#)

| Result<br>No. | Query |       |        |    |          | Description        |
|---------------|-------|-------|--------|----|----------|--------------------|
|               | Score | Match | Length | DB | ID       |                    |
| 1             | 45    | 100.0 | 9      | 2  | AAW99196 | Aaw99196 Minor his |
| 2             | 45    | 100.0 | 9      | 2  | AAW97375 | Aaw97375 HA-1 H-al |
| 3             | 45    | 100.0 | 9      | 8  | ADH40333 | Adh40333 Human min |
| 4             | 37    | 82.2  | 9      | 2  | AAW99197 | Aaw99197 Minor his |
| 5             | 37    | 82.2  | 9      | 2  | AAW97374 | Aaw97374 HA-1 R-al |
| 6             | 37    | 82.2  | 9      | 8  | ADH40334 | Adh40334 Human min |
| 7             | 36    | 80.0  | 9      | 2  | AAW99195 | Aaw99195 Minor his |
| 8             | 36    | 80.0  | 9      | 2  | AAW97572 | Aaw97572 T-cell ep |
| 9             | 31    | 68.9  | 9      | 8  | ADU99853 | Adu99853 BCZ4 tumo |
| 10            | 30    | 66.7  | 9      | 2  | AAW97373 | Aaw97373 Peptide e |
| 11            | 28    | 62.2  | 9      | 2  | AAY10122 | Aay10122 T cell ep |
| 12            | 28    | 62.2  | 9      | 5  | ABG79805 | Abg79805 MHC class |
| 13            | 28    | 62.2  | 9      | 8  | ADK68732 | Adk68732 Epitope l |
| 14            | 28    | 62.2  | 9      | 8  | ADK05291 | Adk05291 Hepatitis |
| 15            | 28    | 62.2  | 9      | 8  | ADK05293 | Adk05293 Hepatitis |
| 16            | 28    | 62.2  | 9      | 8  | ADQ10530 | Adq10530 Cercopith |
| 17            | 28    | 62.2  | 9      | 8  | ADS81010 | Ads81010 Tumour-as |
| 18            | 27    | 60.0  | 9      | 6  | ABJ20115 | Abj20115 MHC bindi |
| 19            | 27    | 60.0  | 9      | 8  | ADT02787 | Adt02787 Human tum |
| 20            | 26    | 57.8  | 6      | 6  | ABU97233 | Abu97233 Enzyme pe |
| 21            | 26    | 57.8  | 6      | 6  | ABU97232 | Abu97232 Enzyme pe |
| 22            | 26    | 57.8  | 6      | 6  | ABU97234 | Abu97234 Enzyme pe |
| 23            | 26    | 57.8  | 6      | 6  | ABU97231 | Abu97231 Enzyme pe |
| 24            | 26    | 57.8  | 6      | 9  | ADY63770 | Ady63770 Human alb |
| 25            | 26    | 57.8  | 7      | 8  | ADM96262 | Adm96262 Human ser |
| 26            | 26    | 57.8  | 9      | 5  | AAE31275 | Aae31275 Human mag |
| 27            | 26    | 57.8  | 9      | 8  | ADM96261 | Adm96261 Human ser |
| 28            | 26    | 57.8  | 9      | 8  | ADM96258 | Adm96258 Human ser |
| 29            | 26    | 57.8  | 9      | 8  | ADT73023 | Adt73023 Human RSV |
| 30            | 26    | 57.8  | 9      | 8  | ADT49074 | Adt49074 Human BFA |
| 31            | 26    | 57.8  | 9      | 8  | ADU99636 | Adu99636 BFA5 tumo |
| 32            | 25    | 55.6  | 6      | 6  | ABU97229 | Abu97229 Enzyme pe |
| 33            | 25    | 55.6  | 6      | 6  | ABU97230 | Abu97230 Enzyme pe |
| 34            | 25    | 55.6  | 9      | 4  | AAG79144 | Aag79144 Amino ter |
| 35            | 25    | 55.6  | 9      | 4  | AAB84492 | Aab84492 Peptide f |
| 36            | 25    | 55.6  | 9      | 5  | AAE26631 | Aae26631 Yeast GPA |
| 37            | 25    | 55.6  | 9      | 7  | ADC25889 | Adc25889 Yeast GPA |
| 38            | 24    | 53.3  | 6      | 2  | AAR29358 | Aar29358 Endotheli |
| 39            | 24    | 53.3  | 6      | 2  | AAR69140 | Aar69140 Endotheli |
| 40            | 24    | 53.3  | 7      | 2  | AAW33386 | Aaw33386 Altered I |
| 41            | 24    | 53.3  | 7      | 4  | AAB70589 | Aab70589 Human imm |
| 42            | 24    | 53.3  | 7      | 5  | AAO21066 | Aao21066 Isomerise |
| 43            | 24    | 53.3  | 7      | 5  | AAE28095 | Aae28095 Human imm |
| 44            | 24    | 53.3  | 8      | 8  | ADT39474 | Adt39474 hSARS vir |
| 45            | 24    | 53.3  | 8      | 8  | ADS78893 | Ads78893 SARS viru |

## ALIGNMENTS

RESULT 1  
 AAW99196  
 ID AAW99196 standard; peptide; 9 AA.  
 XX  
 AC AAW99196;  
 XX  
 DT 20-MAY-1999 (first entry)

# SCORE Search Results Details for Application 10623176 and Search Result us-10-623-176a- 2.rai.

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This page gives you Search Results detail for the Application 10623176 and Search Result us-10-623-176a-2.rai.

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OM protein - protein search, using sw model

Run on: August 18, 2006, 00:49:01 ; Search time 49 Seconds  
(without alignments)  
16.077 Million cell updates/sec

Title: US-10-623-176A-2

Perfect score: 45

Sequence: 1 VLHDDLLEA 9

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 650591 seqs, 87530628 residues

Total number of hits satisfying chosen parameters: 145183

Minimum DB seq length: 0

Maximum DB seq length: 9

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : Issued\_Patents\_AA:  
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2: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/6\_COMB.pep:  
3: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/7\_COMB.pep:  
4: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/H\_COMB.pep:  
5: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/PCTUS\_COMB.pep:  
6: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/RE\_COMB.pep:  
7: /EMC\_Celerra\_SIDS3/ptodata/2/iaa/backfiles1.pep:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

| Result | Query |       |        |    |             |
|--------|-------|-------|--------|----|-------------|
| No.    | Score | Match | Length | DB | ID          |
|        |       |       |        |    | Description |

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| 1  | 45   | 100.0 | 9 | 2 | US-09-269-250E-20    | Sequence 20, Appl |
| 2  | 45   | 100.0 | 9 | 2 | US-09-489-760-2      | Sequence 2, Appl  |
| 3  | 37   | 82.2  | 9 | 2 | US-09-269-250E-18    | Sequence 18, Appl |
| 4  | 37   | 82.2  | 9 | 2 | US-09-489-760-5      | Sequence 5, Appl  |
| 5  | 36   | 80.0  | 9 | 2 | US-09-269-250E-29    | Sequence 29, Appl |
| 6  | 36   | 80.0  | 9 | 2 | US-09-489-760-1      | Sequence 1, Appl  |
| 7  | 30   | 66.7  | 9 | 2 | US-09-489-760-4      | Sequence 4, Appl  |
| 8  | 28.5 | 63.3  | 8 | 2 | US-09-269-250E-38    | Sequence 38, Appl |
| 9  | 28   | 62.2  | 9 | 2 | US-09-776-232-52     | Sequence 52, Appl |
| 10 | 28   | 62.2  | 9 | 2 | US-09-380-534-52     | Sequence 52, Appl |
| 11 | 26   | 57.8  | 6 | 2 | US-10-166-225A-163   | Sequence 163, App |
| 12 | 26   | 57.8  | 6 | 2 | US-10-166-225A-164   | Sequence 164, App |
| 13 | 26   | 57.8  | 6 | 2 | US-10-166-225A-165   | Sequence 165, App |
| 14 | 26   | 57.8  | 6 | 2 | US-10-166-225A-166   | Sequence 166, App |
| 15 | 25   | 55.6  | 6 | 2 | US-10-166-225A-161   | Sequence 161, App |
| 16 | 25   | 55.6  | 6 | 2 | US-10-166-225A-162   | Sequence 162, App |
| 17 | 25   | 55.6  | 7 | 2 | US-09-025-819-5      | Sequence 5, Appl  |
| 18 | 25   | 55.6  | 7 | 2 | US-09-808-126-5      | Sequence 5, Appl  |
| 19 | 25   | 55.6  | 7 | 2 | US-09-803-951-5      | Sequence 5, Appl  |
| 20 | 25   | 55.6  | 9 | 2 | US-08-582-333A-3     | Sequence 3, Appl  |
| 21 | 25   | 55.6  | 9 | 2 | US-09-305-923A-7     | Sequence 7, Appl  |
| 22 | 25   | 55.6  | 9 | 2 | US-08-946-298-4      | Sequence 4, Appl  |
| 23 | 24   | 53.3  | 9 | 2 | US-09-776-232-491    | Sequence 491, App |
| 24 | 24   | 53.3  | 9 | 2 | US-09-380-534-491    | Sequence 491, App |
| 25 | 24   | 53.3  | 9 | 3 | US-09-641-528B-18057 | Sequence 18057, A |
| 26 | 24   | 53.3  | 9 | 3 | US-09-641-528B-48631 | Sequence 48631, A |
| 27 | 23   | 51.1  | 6 | 2 | US-09-217-609A-8     | Sequence 8, Appl  |
| 28 | 23   | 51.1  | 6 | 2 | US-08-873-235B-8     | Sequence 8, Appl  |
| 29 | 23   | 51.1  | 9 | 2 | US-08-159-339A-370   | Sequence 370, App |
| 30 | 22   | 48.9  | 5 | 2 | US-08-811-463-30     | Sequence 30, Appl |
| 31 | 22   | 48.9  | 5 | 2 | US-09-933-497B-30    | Sequence 30, Appl |
| 32 | 22   | 48.9  | 7 | 1 | US-08-208-036-7      | Sequence 7, Appl  |
| 33 | 22   | 48.9  | 7 | 1 | US-08-208-036-9      | Sequence 9, Appl  |
| 34 | 22   | 48.9  | 7 | 1 | US-08-428-823-7      | Sequence 7, Appl  |
| 35 | 22   | 48.9  | 7 | 1 | US-08-428-823-9      | Sequence 9, Appl  |
| 36 | 22   | 48.9  | 7 | 2 | US-08-556-419-14     | Sequence 14, Appl |
| 37 | 22   | 48.9  | 7 | 2 | US-09-173-941-82     | Sequence 82, Appl |
| 38 | 22   | 48.9  | 7 | 2 | US-09-494-190-82     | Sequence 82, Appl |
| 39 | 22   | 48.9  | 8 | 1 | US-08-403-378B-13    | Sequence 13, Appl |
| 40 | 22   | 48.9  | 9 | 2 | US-09-217-609A-11    | Sequence 11, Appl |
| 41 | 22   | 48.9  | 9 | 2 | US-08-873-235B-11    | Sequence 11, Appl |
| 42 | 22   | 48.9  | 9 | 3 | US-09-641-528B-32008 | Sequence 32008, A |
| 43 | 22   | 48.9  | 9 | 3 | US-09-641-528B-40149 | Sequence 40149, A |
| 44 | 21   | 46.7  | 6 | 1 | US-08-459-568-20     | Sequence 20, Appl |
| 45 | 21   | 46.7  | 6 | 1 | US-08-399-411-20     | Sequence 20, Appl |

## ALIGNMENTS

## RESULT 1

US-09-269-250E-20

; Sequence 20, Application US/09269250E  
; Patent No. 6830883  
; GENERAL INFORMATION:  
; APPLICANT: Goulmy, Elsa  
; TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1  
; FILE REFERENCE: 58994  
; CURRENT APPLICATION NUMBER: US/09/269,250E  
; CURRENT FILING DATE: 1999-05-21

# SCORE Search Results Details for Application 10623176 and Search Result us-10-623-176a 2.rapbm.

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OM protein - protein search, using sw model

Run on: August 18, 2006, 01:00:36 ; Search time 179 Seconds  
(without alignments)  
23.290 Million cell updates/sec

Title: US-10-623-176A-2

Perfect score: 45

Sequence: 1 VLHDDLLEA 9

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 2097797 seqs, 463214858 residues

Total number of hits satisfying chosen parameters: 222252

Minimum DB seq length: 0

Maximum DB seq length: 9

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : Published\_Applications\_AA\_Main:  
1: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US07\_PUBCOMB.pep:  
2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US08\_PUBCOMB.pep:  
3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US09\_PUBCOMB.pep:  
4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10A\_PUBCOMB.pep:  
5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10B\_PUBCOMB.pep:  
6: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US11\_PUBCOMB.pep:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

| Result<br>No. | Query<br>Score | %<br>Match Length DB ID |   |                 |                 | Description       |
|---------------|----------------|-------------------------|---|-----------------|-----------------|-------------------|
|               |                | 9                       | 4 | US-10-623-176-2 |                 |                   |
| 1             | 45             | 100.0                   | 9 | 4               | US-10-623-176-2 | Sequence 2, Appli |

|    |      |       |   |   |                    |                   |
|----|------|-------|---|---|--------------------|-------------------|
| 2  | 45   | 100.0 | 9 | 4 | US-10-791-217-2    | Sequence 2, Appli |
| 3  | 45   | 100.0 | 9 | 5 | US-10-861-335-1    | Sequence 1, Appli |
| 4  | 45   | 100.0 | 9 | 6 | US-11-007-740-20   | Sequence 20, Appl |
| 5  | 45   | 100.0 | 9 | 6 | US-11-010-748A-11  | Sequence 11, Appl |
| 6  | 41   | 91.1  | 9 | 4 | US-10-623-176-41   | Sequence 41, Appl |
| 7  | 41   | 91.1  | 9 | 4 | US-10-623-176-45   | Sequence 45, Appl |
| 8  | 37   | 82.2  | 9 | 4 | US-10-623-176-10   | Sequence 10, Appl |
| 9  | 37   | 82.2  | 9 | 4 | US-10-623-176-47   | Sequence 47, Appl |
| 10 | 37   | 82.2  | 9 | 4 | US-10-791-217-5    | Sequence 5, Appli |
| 11 | 37   | 82.2  | 9 | 6 | US-11-007-740-18   | Sequence 18, Appl |
| 12 | 37   | 82.2  | 9 | 6 | US-11-010-748A-12  | Sequence 12, Appl |
| 13 | 36   | 80.0  | 9 | 4 | US-10-623-176-1    | Sequence 1, Appli |
| 14 | 36   | 80.0  | 9 | 4 | US-10-623-176-40   | Sequence 40, Appl |
| 15 | 36   | 80.0  | 9 | 4 | US-10-791-217-1    | Sequence 1, Appli |
| 16 | 36   | 80.0  | 9 | 6 | US-11-007-740-29   | Sequence 29, Appl |
| 17 | 33   | 73.3  | 9 | 4 | US-10-623-176-42   | Sequence 42, Appl |
| 18 | 33   | 73.3  | 9 | 4 | US-10-623-176-46   | Sequence 46, Appl |
| 19 | 32   | 71.1  | 9 | 4 | US-10-623-176-4    | Sequence 4, Appli |
| 20 | 30   | 66.7  | 9 | 4 | US-10-623-176-14   | Sequence 14, Appl |
| 21 | 30   | 66.7  | 9 | 4 | US-10-791-217-4    | Sequence 4, Appli |
| 22 | 30   | 66.7  | 9 | 6 | US-11-007-740-40   | Sequence 40, Appl |
| 23 | 29   | 64.4  | 9 | 4 | US-10-623-176-48   | Sequence 48, Appl |
| 24 | 28.5 | 63.3  | 8 | 4 | US-10-623-176-15   | Sequence 15, Appl |
| 25 | 28.5 | 63.3  | 8 | 6 | US-11-007-740-38   | Sequence 38, Appl |
| 26 | 28   | 62.2  | 9 | 4 | US-10-623-176-23   | Sequence 23, Appl |
| 27 | 28   | 62.2  | 9 | 4 | US-10-623-176-38   | Sequence 38, Appl |
| 28 | 28   | 62.2  | 9 | 4 | US-10-777-053-95   | Sequence 95, Appl |
| 29 | 28   | 62.2  | 9 | 4 | US-10-837-217-95   | Sequence 95, Appl |
| 30 | 27   | 60.0  | 9 | 5 | US-10-705-459-280  | Sequence 280, App |
| 31 | 26   | 57.8  | 6 | 4 | US-10-166-225A-163 | Sequence 163, App |
| 32 | 26   | 57.8  | 6 | 4 | US-10-166-225A-164 | Sequence 164, App |
| 33 | 26   | 57.8  | 6 | 4 | US-10-166-225A-165 | Sequence 165, App |
| 34 | 26   | 57.8  | 6 | 4 | US-10-166-225A-166 | Sequence 166, App |
| 35 | 26   | 57.8  | 6 | 6 | US-11-129-143-163  | Sequence 163, App |
| 36 | 26   | 57.8  | 6 | 6 | US-11-129-143-164  | Sequence 164, App |
| 37 | 26   | 57.8  | 6 | 6 | US-11-129-143-165  | Sequence 165, App |
| 38 | 26   | 57.8  | 6 | 6 | US-11-129-143-166  | Sequence 166, App |
| 39 | 26   | 57.8  | 9 | 5 | US-10-825-026-48   | Sequence 48, Appl |
| 40 | 25   | 55.6  | 6 | 4 | US-10-166-225A-161 | Sequence 161, App |
| 41 | 25   | 55.6  | 6 | 4 | US-10-166-225A-162 | Sequence 162, App |
| 42 | 25   | 55.6  | 6 | 6 | US-11-129-143-161  | Sequence 161, App |
| 43 | 25   | 55.6  | 6 | 6 | US-11-129-143-162  | Sequence 162, App |
| 44 | 25   | 55.6  | 7 | 3 | US-09-803-951-5    | Sequence 5, Appli |
| 45 | 25   | 55.6  | 9 | 3 | US-09-747-774A-3   | Sequence 3, Appli |

## ALIGNMENTS

## RESULT 1

US-10-623-176-2

```
; Sequence 2, Application US/10623176
; Publication No. US20040092446A1
; GENERAL INFORMATION:
; APPLICANT: Goulimy, Els A.J.M.
; APPLICANT: Hunt, Donald F.
; APPLICANT: Engelhard, Victor H.
; TITLE OF INVENTION: HA-1 epitopes and uses thereof
; FILE REFERENCE: 2183-6047US
; CURRENT APPLICATION NUMBER: US/10/623,176
; CURRENT FILING DATE: 2003-07-18
```

# SCORE Search Results Details for Application 10623176 and Search Result us-10-623-176a- 2.rapbn.

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|                                 |   |                                       |                           |  |
|---------------------------------|---|---------------------------------------|---------------------------|--|
| <a href="#">Score Home Page</a> | <a href="#">Retrieve Application List</a> | <a href="#">SCORE System Overview</a> | <a href="#">SCORE FAQ</a> | <a href="#">Comments / Suggestions</a> |
|---------------------------------|---|---------------------------------------|---------------------------|--|

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This page gives you Search Results detail for the Application 10623176 and Search Result us-10-623-176a-2.rapbn.

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GenCore version 5.1.9  
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OM protein - protein search, using sw model

Run on: August 18, 2006, 01:01:12 ; Search time 32 Seconds  
(without alignments)  
19.056 Million cell updates/sec

Title: US-10-623-176A-2

Perfect score: 45

Sequence: 1 VLHDDLLEA 9

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 243793 seqs, 67754213 residues

Total number of hits satisfying chosen parameters: 13145

Minimum DB seq length: 0

Maximum DB seq length: 9

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : Published\_Applications\_AA\_New:  
1: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US09\_NEW\_PUB.pep:  
2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US06\_NEW\_PUB.pep:  
3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US07\_NEW\_PUB.pep:  
4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US08\_NEW\_PUB.pep:  
5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/PCT\_NEW\_PUB.pep:  
6: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US10\_NEW\_PUB.pep:  
7: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US11\_NEW\_PUB.pep:  
8: /EMC\_Celerra\_SIDS3/ptodata/2/pubpaa/US60\_NEW\_PUB.pep:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

|        |   |       |
|--------|---|-------|
| Result | % | Query |
|--------|---|-------|

| No. | Score | Match Length | DB | ID                    | Description       |
|-----|-------|--------------|----|-----------------------|-------------------|
| 1   | 28    | 62.2         | 9  | 7 US-11-313-152-52    | Sequence 52, Appl |
| 2   | 24    | 53.3         | 7  | 7 US-11-263-230-344   | Sequence 344, App |
| 3   | 24    | 53.3         | 9  | 7 US-11-313-152-491   | Sequence 491, App |
| 4   | 23    | 51.1         | 9  | 6 US-10-537-642-611   | Sequence 611, App |
| 5   | 22    | 48.9         | 8  | 7 US-11-122-986-753   | Sequence 753, App |
| 6   | 22    | 48.9         | 9  | 6 US-10-781-659-25    | Sequence 25, Appl |
| 7   | 21    | 46.7         | 9  | 7 US-11-140-487A-1758 | Sequence 1758, Ap |
| 8   | 20    | 44.4         | 4  | 6 US-10-514-263-18    | Sequence 18, Appl |
| 9   | 20    | 44.4         | 4  | 6 US-10-511-436A-121  | Sequence 121, App |
| 10  | 20    | 44.4         | 4  | 7 US-11-324-947-86    | Sequence 86, Appl |
| 11  | 20    | 44.4         | 4  | 7 US-11-053-045-2     | Sequence 2, Appli |
| 12  | 20    | 44.4         | 4  | 7 US-11-271-235-5     | Sequence 5, Appli |
| 13  | 20    | 44.4         | 4  | 7 US-11-249-061-5     | Sequence 5, Appli |
| 14  | 20    | 44.4         | 7  | 6 US-10-479-225A-19   | Sequence 19, Appl |
| 15  | 20    | 44.4         | 7  | 6 US-10-479-225A-20   | Sequence 20, Appl |
| 16  | 20    | 44.4         | 7  | 7 US-11-398-130-8     | Sequence 8, Appli |
| 17  | 19    | 42.2         | 7  | 7 US-11-122-986-362   | Sequence 362, App |
| 18  | 19    | 42.2         | 7  | 7 US-11-263-230-1202  | Sequence 1202, Ap |
| 19  | 19    | 42.2         | 7  | 7 US-11-263-230-1486  | Sequence 1486, Ap |
| 20  | 19    | 42.2         | 7  | 7 US-11-263-230-1487  | Sequence 1487, Ap |
| 21  | 19    | 42.2         | 7  | 7 US-11-398-130-16    | Sequence 16, Appl |
| 22  | 19    | 42.2         | 9  | 6 US-10-537-642-323   | Sequence 323, App |
| 23  | 19    | 42.2         | 9  | 6 US-10-570-010-25    | Sequence 25, Appl |
| 24  | 19    | 42.2         | 9  | 7 US-11-140-487A-1828 | Sequence 1828, Ap |
| 25  | 18    | 40.0         | 7  | 7 US-11-263-230-362   | Sequence 362, App |
| 26  | 18    | 40.0         | 7  | 7 US-11-263-230-651   | Sequence 651, App |
| 27  | 18    | 40.0         | 7  | 7 US-11-263-230-790   | Sequence 790, App |
| 28  | 18    | 40.0         | 7  | 7 US-11-263-230-934   | Sequence 934, App |
| 29  | 18    | 40.0         | 7  | 7 US-11-263-230-1067  | Sequence 1067, Ap |
| 30  | 18    | 40.0         | 7  | 7 US-11-263-230-1346  | Sequence 1346, Ap |
| 31  | 18    | 40.0         | 8  | 7 US-11-223-610-23    | Sequence 23, Appl |
| 32  | 18    | 40.0         | 8  | 7 US-11-234-731-622   | Sequence 622, App |
| 33  | 18    | 40.0         | 9  | 6 US-10-781-659-60    | Sequence 60, Appl |
| 34  | 18    | 40.0         | 9  | 6 US-10-537-642-471   | Sequence 471, App |
| 35  | 18    | 40.0         | 9  | 7 US-11-140-487A-1073 | Sequence 1073, Ap |
| 36  | 18    | 40.0         | 9  | 7 US-11-140-487A-1227 | Sequence 1227, Ap |
| 37  | 18    | 40.0         | 9  | 7 US-11-122-986-817   | Sequence 817, App |
| 38  | 18    | 40.0         | 9  | 7 US-11-240-651-10    | Sequence 10, Appl |
| 39  | 18    | 40.0         | 9  | 7 US-11-011-026-24    | Sequence 24, Appl |
| 40  | 18    | 40.0         | 9  | 7 US-11-313-152-155   | Sequence 155, App |
| 41  | 18    | 40.0         | 9  | 7 US-11-313-152-417   | Sequence 417, App |
| 42  | 17    | 37.8         | 6  | 7 US-11-178-155-13    | Sequence 13, Appl |
| 43  | 17    | 37.8         | 6  | 7 US-11-287-157A-163  | Sequence 163, App |
| 44  | 17    | 37.8         | 7  | 1 US-09-906-481E-7    | Sequence 7, Appli |
| 45  | 17    | 37.8         | 7  | 7 US-11-023-959A-29   | Sequence 29, Appl |

## ALIGNMENTS

## RESULT 1

US-11-313-152-52

; Sequence 52, Application US/11313152  
; Publication No. US20060153858A1  
; GENERAL INFORMATION:  
; APPLICANT: Kundig, Thomas M.  
; APPLICANT: Simard, John J. L.  
; TITLE OF INVENTION: METHOD OF INDUCING A CTL RESPONSE  
; FILE REFERENCE: MANNK.001CP2C1

# SCORE Search Results Details

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This page gives you Search Results detail for the Application 10623176 and Search Result us-10-6; start

GenCore version 5.1.9  
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OM protein - protein search, using sw model

Run on: August 18, 2006, 00:44:01 ; Search time 38 Seconds  
(without alignments)  
22.788 Million cell updates/sec

Title: US-10-623-176A-2

Perfect score: 45

Sequence: 1 VLHDDLLEA 9

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 791

Minimum DB seq length: 0

Maximum DB seq length: 9

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : PIR\_80:\*

1: pir1:\*

2: pir2:\*

3: pir3:\*

4: pir4:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

%

| Result No. | Query |       |        |       | Description |
|------------|-------|-------|--------|-------|-------------|
|            | Score | Match | Length | DB ID |             |

|   |    |      |   |   |        |                    |
|---|----|------|---|---|--------|--------------------|
| 1 | 18 | 40.0 | 7 | 2 | I46868 | alpha-myosin heavy |
| 2 | 17 | 37.8 | 9 | 2 | S55696 | phosphoenolpyruvat |
| 3 | 16 | 35.6 | 7 | 2 | PQ0663 | membrane protein - |
| 4 | 16 | 35.6 | 7 | 2 | S68004 | hucolin, 75K chain |
| 5 | 16 | 35.6 | 8 | 2 | PC4131 | hypothetical prote |
| 6 | 15 | 33.3 | 6 | 2 | S78764 | ribosomal protein  |
| 7 | 15 | 33.3 | 7 | 2 | A59489 | protein kinase C i |
| 8 | 14 | 31.1 | 7 | 2 | S20446 | elastase - Pseudom |

|    |    |      |   |   |        |                    |
|----|----|------|---|---|--------|--------------------|
| 9  | 14 | 31.1 | 8 | 2 | XGHUEU | urine glycopeptide |
| 10 | 14 | 31.1 | 9 | 2 | A12872 | transaldolase (EC  |
| 11 | 14 | 31.1 | 9 | 2 | S10920 | venom protein HR-3 |
| 12 | 14 | 31.1 | 9 | 2 | A60427 | macrophage cytotox |
| 13 | 13 | 28.9 | 8 | 2 | A61328 | trypsin (EC 3.4.21 |
| 14 | 13 | 28.9 | 9 | 2 | PH0942 | T-cell receptor be |
| 15 | 12 | 26.7 | 4 | 2 | I40697 | biotin A - Citroba |
| 16 | 12 | 26.7 | 5 | 2 | PT0679 | T-cell receptor be |
| 17 | 12 | 26.7 | 5 | 2 | PT0601 | T-cell receptor be |
| 18 | 12 | 26.7 | 6 | 2 | B35640 | cerebellar degener |
| 19 | 12 | 26.7 | 6 | 2 | PT0533 | T-cell receptor be |
| 20 | 12 | 26.7 | 7 | 2 | A34026 | acetylcholinestera |
| 21 | 12 | 26.7 | 7 | 2 | B39040 | calsequestrin, fas |
| 22 | 12 | 26.7 | 7 | 2 | PT0628 | T-cell receptor be |
| 23 | 12 | 26.7 | 7 | 2 | PT0722 | T-cell receptor be |
| 24 | 12 | 26.7 | 7 | 2 | PT0576 | T-cell receptor be |
| 25 | 12 | 26.7 | 8 | 2 | PT0368 | Ig gamma chain C r |
| 26 | 12 | 26.7 | 8 | 2 | PN0043 | phosphatidylethano |
| 27 | 12 | 26.7 | 8 | 2 | PT0557 | T-cell receptor be |
| 28 | 12 | 26.7 | 9 | 2 | A60108 | exotoxin A - Strep |
| 29 | 12 | 26.7 | 9 | 2 | PW0002 | chlorophyll a/b-bi |
| 30 | 12 | 26.7 | 9 | 2 | S65913 | pyrimidine synthes |
| 31 | 12 | 26.7 | 9 | 2 | PH0108 | late G1-69 protein |
| 32 | 12 | 26.7 | 9 | 2 | PT0562 | T-cell receptor be |
| 33 | 12 | 26.7 | 9 | 2 | B30572 | T-cell receptor be |
| 34 | 11 | 24.4 | 5 | 2 | C41225 | copper resistance  |
| 35 | 11 | 24.4 | 5 | 2 | T10954 | hypothetical prote |
| 36 | 11 | 24.4 | 6 | 2 | T11779 | phosphoglycerate t |
| 37 | 11 | 24.4 | 7 | 2 | S25266 | pilE protein - Esc |
| 38 | 11 | 24.4 | 7 | 2 | PT0246 | Ig heavy chain CRD |
| 39 | 11 | 24.4 | 8 | 2 | S22428 | chitin-binding pro |
| 40 | 11 | 24.4 | 8 | 2 | B33099 | 158K exoantigen -  |
| 41 | 11 | 24.4 | 8 | 2 | S69165 | ferredoxin a2 - Ja |
| 42 | 11 | 24.4 | 9 | 2 | S66419 | tetrameric protein |
| 43 | 11 | 24.4 | 9 | 2 | PT0272 | Ig heavy chain CRD |
| 44 | 11 | 24.4 | 9 | 2 | A11497 | transaldolase (EC  |
| 45 | 11 | 24.4 | 9 | 2 | B39504 | octamer-binding pr |

## ALIGNMENTS

## RESULT 1

I46868

alpha-myosin heavy chain - rabbit (fragment)

C;Species: Oryctolagus cuniculus (domestic rabbit)

C;Date: 14-Feb-1997 #sequence\_revision 14-Feb-1997 #text\_change 09-Jul-2004

C;Accession: I46868

R;Friedman, D.J.; Umeda, P.K.; Sinha, A.M.; Hsu, H.

Proc. Natl. Acad. Sci. U.S.A. 81, 3044-3048, 1984

A;Title: Characterization of genomic clones specifying rabbit alpha- and beta-ventricu

A;Reference number: I46868; MUID:84221901; PMID:6328491

A;Accession: I46868

A;Status: preliminary; translated from GB/EMBL/DDBJ

A;Molecule type: DNA

A;Residues: 1-7

A;Cross-references: UNIPROT:Q28742; UNIPARC:UPI0000087938; GB:K01698; NID:g165538; PID

Query Match 40.0%; Score 18; DB 2; Length 7;

Best Local Similarity 50.0%; Pred. No. 2.8e+05;

Matches 2; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

# **SCORE Search Results Details for Application 10623176 and Search Result us-10-623-176a- 2.rup.**

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This page gives you Search Results detail for the Application 10623176 and Search Result us-10-623-176a-2.rup.

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## OM protein - protein search, using sw model

Run on: August 18, 2006, 00:40:41 ; Search time 295 Seconds  
(without alignments)  
28.221 Million cell updates/sec

Title: US-10-623-176A-2

Perfect score: 45

Sequence: 1 VLHDDLLEA 9

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2849598 seqs, 925015592 residues

Total number of hits satisfying chosen parameters: 2195

Minimum DB seq length: 0

Maximum DB seq length: 9

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : UniProt\_7.2:\*  
1: uniprot\_sprot:/\*  
2: uniprot\_trembl:/\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

8

| Result<br>No. | Score | Query |        |    |               |  | Description        |
|---------------|-------|-------|--------|----|---------------|--|--------------------|
|               |       | Match | Length | DB | ID            |  |                    |
| 1             | 19    | 42.2  | 9      | 2  | Q7R8X5_PLAYO  |  | Q7r8x5 plasmodium  |
| 2             | 19    | 42.2  | 9      | 2  | Q47556_ECOLI  |  | Q47556 escherichia |
| 3             | 18    | 40.0  | 7      | 2  | Q28742_RABBIT |  | Q28742 oryctolagus |
| 4             | 17    | 37.8  | 9      | 2  | Q7M3S5_9TRYP  |  | Q7m3s5 trypanosoma |

|    |    |      |   |   |               |                     |
|----|----|------|---|---|---------------|---------------------|
| 5  | 16 | 35.6 | 7 | 2 | O99182_9SMEG  | O99182 gnatholebia  |
| 6  | 16 | 35.6 | 8 | 2 | Q7Z6G0_HUMAN  | Q7z6g0 homo sapien  |
| 7  | 16 | 35.6 | 8 | 2 | Q2L8A3_COFC   | Q218a3 coffeea cane |
| 8  | 16 | 35.6 | 8 | 2 | Q2L8A4_9GENT  | Q218a4 coffeea cong |
| 9  | 16 | 35.6 | 9 | 1 | FAR8_MACRS    | P83281 macrobrachi  |
| 10 | 16 | 35.6 | 9 | 2 | Q70SM2_HUMAN  | Q70sm2 homo sapien  |
| 11 | 16 | 35.6 | 9 | 2 | Q2Q2A8_CAEEL  | Q2q2a8 caenorhabdi  |
| 12 | 16 | 35.6 | 9 | 2 | Q4X981_PLACH  | Q4x981 plasmodium   |
| 13 | 16 | 35.6 | 9 | 2 | Q8LPT5_MAIZE  | Q8lpt5 zea mays (m  |
| 14 | 16 | 35.6 | 9 | 2 | P82568_STRPY  | P82568 streptococc  |
| 15 | 16 | 35.6 | 9 | 2 | Q2WDC8_FICHY  | Q2wdc8 ficedula hy  |
| 16 | 16 | 35.6 | 9 | 2 | Q2WDC9_FICPA  | Q2wdc9 ficedula pa  |
| 17 | 16 | 35.6 | 9 | 2 | Q2WDD7_FICAL  | Q2wdd7 ficedula al  |
| 18 | 15 | 33.3 | 8 | 2 | Q15900_HUMAN  | Q15900 homo sapien  |
| 19 | 15 | 33.3 | 8 | 2 | Q4XT27_PLACH  | Q4xt27 plasmodium   |
| 20 | 15 | 33.3 | 9 | 2 | Q9XJN0_9VIRU  | Q9xjn0 bacteriopha  |
| 21 | 14 | 31.1 | 6 | 1 | CWP29_ARATH   | P80849 arabidopsis  |
| 22 | 14 | 31.1 | 6 | 1 | TRPI_PSEPU    | P36414 pseudomonas  |
| 23 | 14 | 31.1 | 8 | 1 | GLUR_HUMAN    | P02729 homo sapien  |
| 24 | 14 | 31.1 | 8 | 2 | Q9BFA7_MACPR  | Q9bfa7 macroscelid  |
| 25 | 14 | 31.1 | 8 | 2 | P72279_RHOGO  | P72279 rhodococcus  |
| 26 | 14 | 31.1 | 8 | 2 | Q93SR0_STAEP  | Q93sr0 staphylococ  |
| 27 | 14 | 31.1 | 8 | 2 | Q3S8H3_9HIV1  | Q3s8h3 human immun  |
| 28 | 14 | 31.1 | 9 | 1 | TAL1_PICJA    | P17440 pichia jadi  |
| 29 | 14 | 31.1 | 9 | 2 | Q7M4R5_HUMAN  | Q7m4r5 homo sapien  |
| 30 | 14 | 31.1 | 9 | 2 | Q7M471_VESOR  | Q7m471 vespa orien  |
| 31 | 14 | 31.1 | 9 | 2 | Q9FSZ2_CICAR  | Q9fsz2 cicer ariet  |
| 32 | 14 | 31.1 | 9 | 2 | Q8CG39_RAT    | Q8cg39 rattus norv  |
| 33 | 14 | 31.1 | 9 | 2 | Q9QZA8_MOUSE  | Q9qza8 mus musculu  |
| 34 | 14 | 31.1 | 9 | 2 | Q6Q7G0_RANRI  | Q6q7g0 rana ridibu  |
| 35 | 14 | 31.1 | 9 | 2 | Q85710_9RETR  | Q85710 rous sarcom  |
| 36 | 14 | 31.1 | 9 | 2 | Q8UTD7_9HIV1  | Q8utd7 human immun  |
| 37 | 13 | 28.9 | 7 | 2 | Q66205_9CORS  | Q66205 transmissib  |
| 38 | 13 | 28.9 | 8 | 2 | Q9HDS4_ASPLFL | Q9hds4 aspergillus  |
| 39 | 13 | 28.9 | 8 | 2 | Q15889_HUMAN  | Q15889 homo sapien  |
| 40 | 13 | 28.9 | 8 | 2 | Q7M390_BALAC  | Q7m390 balaenopter  |
| 41 | 13 | 28.9 | 8 | 2 | Q37854_BPR17  | Q37854 bacteriopha  |
| 42 | 13 | 28.9 | 8 | 2 | Q6JC68_SOYBN  | Q6jc68 glycine max  |
| 43 | 13 | 28.9 | 8 | 2 | Q7XB03_MAIZE  | Q7xb03 zea mays (m  |
| 44 | 13 | 28.9 | 8 | 2 | Q51594_9ZZZZ  | Q51594 plasmid col  |
| 45 | 13 | 28.9 | 9 | 2 | Q56SS9_SAMCA  | Q56ss9 sambucus ca  |

## ALIGNMENTS

RESULT 1  
 Q7R8X5\_PLAYO  
 ID Q7R8X5\_PLAYO PRELIMINARY; PRT; 9 AA.  
 AC Q7R8X5;  
 DT 15-DEC-2003, integrated into UniProtKB/TrEMBL.  
 DT 15-DEC-2003, sequence version 1.  
 DT 07-FEB-2006, entry version 10.  
 DE Hypothetical protein.  
 GN ORFNames=PY07095;  
 OS Plasmodium yoelii yoelii.  
 OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.  
 OX NCBI\_TaxID=73239;  
 RN [1]  
 RP NUCLEOTIDE SEQUENCE [LARGE SCALE GENOMIC DNA].  
 RC STRAIN=17XNL;